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(54) Title: NOVEL CACAO ENDOPROTEINASES AND THEIR USE IN THE PRODUCTION OF COCOA FLAVOUR

(57) Abstract: The present invention pertains to novel aspartic endoproteinases from *Th. cacao* which are involved in the production of cocoa flavour and DNA sequences coding for them. In particular, the present invention relates to the use of said enzymes in the manufacture of cocoa flavour.

**Novel cacao endoproteinases and their use
in the production of cocoa flavour**

5

The present invention pertains to novel endoproteinases involved in the production of cocoa flavour and the DNA coding for them. In particular, the present invention relates to the use of said enzymes for the manufacture of cocoa flavour.

10 It is known that in processing cacao beans the generation of the typical cocoa flavour requires two steps - the fermentation step, which includes air-drying of the fermented material and the roasting step. Though roasting seems to be the key stage of obtaining cocoa flavour subjecting non fermented beans to a roasting step does not yield cocoa flavour suggesting that during the fermentation step precursors are produced that are essential for flavour 15 generation (Rohan J. Food Sci. 29 (1964), 456-459).

During fermentation two major activities may be observed. First, the pulp surrounding the beans is degraded by micro-organisms with the sugars contained in the pulp being largely transformed to acids, especially acetic acid (Quesnel et al. J. Sci. Food. Agric. 16 (1965), 20 441-447; Ostovar and Keeney, J. Food. Sci. 39 (1973), 611-617). The acids then slowly diffuse into the beans and eventually cause an acidification of the cellular material. Second, fermentation also results in a release of peptides exhibiting differing sizes and a generation of a high level of hydrophobic free amino acids. This latter finding led to the hypothesis that proteolysis occurring during the fermentation step is not due to a random protein hydrolysis 25 but seems to be rather based on the activity of specific endoproteinase (Kirchhoff et al., Food Chem 31 (1989), 295-311). This specific mixture of peptides and hydrophobic amino acids is deemed to represent cocoa-specific flavour precursors.

So far in cacao beans several proteolytic enzyme activities have been investigated and

checked for their putative role in the formation of cocoa flavour precursors.

An aspartic endoproteinase activity which is optimal at very low pH (pH 3.5) and is inhibited by pepstatin A has been identified. A polypeptide described to have this activity has been

5 isolated and is described to consist of two peptides (29 and 13 kDa) which are deemed to be derived by self-digestion from a 42 kDa pro-peptide (Voigt et al., J. Plant Physiol. 145 (1995), 299-307). The enzyme cleaves protein substrates between hydrophobic amino acid residues to produce oligopeptides with hydrophobic amino acid residues at the ends (Voigt et al., Food Chem. 49 (1994), 173-180). The enzyme accumulates with the vicilin-class (7S) 10 globulin during bean ripening. Throughout germination, its activity remains constant during the first days and does not decrease before the onset of globulin degradation (Voigt et al., J. Plant Physiol. 145 (1995), 299-307).

A cysteine endoproteinase activity had been isolated which is optimal at a pH of 5. This

15 enzymatic activity is believed not to split native storage proteins in ungerminated seeds. Cysteine endoproteinase activity increases during the germination process when degradation of globular storage protein occurs (Bichl et al., Cocoa Research Conference, Salvador, Bahia, Brasil, 17-23 Nov. 1996).

20 Moreover, a carboxypeptidase activity has been identified which is inhibited by PMSF and thus belongs to the class of serine proteases. It is stable over a broad pH range with a maximum activity at pH 5.8. This enzyme does not degrade native proteins but preferentially splits hydrophobic amino acids from the carboxy-terminus of peptides. Yet, peptides with carboxy-terminal arginine, lysine, or proline residues are seemingly resistant to degradation. 25 The rate of hydrolysis has been found to be not only determined by the carboxy-terminal amino acid as such, but also to be affected by the neighbouring amino acid residue (Bytof et al., Food Chem. 54 (1995), 15-21).

During the second step of cocoa flavour production - the roasting step - the oligopeptides and

30 amino acids generated at the stage of fermentation have been found to obviously undergo a Maillard reaction with reducing sugars present eventually producing the substances

responsible for the cocoa flavour as such. This hypothesis has been confirmed in an experiment, wherein an oligopeptide fraction isolated after fermentation of cacao beans had been subjected to roasting in the presence of free amino acids and reducing sugars to obtain cocoa flavour (Mohr et al., Fette, Seifen, Anstrichmittel 73 (1971), 515-521 and 78 (1976), 88-95).

5

Cocoa-specific aroma has also been obtained in an experiment wherein acetone dry powder (AcDP) prepared from unfermented ripe cacao beans was subjected to autolysis at a pH of 5.2 followed by roasting in the presence of reducing sugars. It was conceived that under these conditions preferentially free hydrophobic amino acids and hydrophilic peptides should be 10 generated and the peptide pattern thus obtained was similar to that of extracts from fermented cacao beans. An analysis of free amino acids revealed that Leu, Ala, Phe and Val were the predominant amino acids liberated in fermented beans or autolysis (Voigt et al., Food Chem. 49 (1994), 173-180). In contrast to these findings no cocoa-specific flavour could be detected when AcDP was subjected to autolysis at a pH of as low as 3.5, the pH, at which the known 15 aspartic endoprotease shows activity. Only few free amino acids were found to be released but a large number of hydrophobic peptides were formed. This may be explained by the aspartic endoprotease having a high activity at this pH with the carboxypeptidase being substantially inactive under these conditions. When incubating peptides obtained after autolysis of AcDP at a pH of 3.5 with carboxypeptidase A from porcine pancreas at pH 7.5 20 hydrophobic amino acids were preferentially released. The pattern of free amino acids and peptides was rather similar to that found in fermented cacao beans and in the proteolysis product obtained by autolysis of AcDP at pH 5.2. After roasting of the amino acids and peptides mixture as above, a cocoa aroma could be generated. On the contrary, with a synthetic mixture of free amino acids alone whose composition was similar to the spectrum 25 found in fermented beans cocoa flavour could not be detected after roasting, indicating that both the peptides and the amino acids are important for this purpose (Voigt et al., Food Chem. 49 (1994), 173-180).

Apart from the enzymes also the protein source of the peptides/amino acids seems to be of 30 importance for the generation of cocoa flavour.

During cacao bean fermentation, the percentage reduction of protein concentration observed for vicilin and albumin was 88.8% and 47.4%, respectively (Amin et al., J. Sci. Food Agric. 76 (1998), 123-128). When peptides obtained by proteolysis of the globulin fraction were post-treated with carboxypeptidase, hydrophobic amino acids (Leu, Phe, Ala, Val, Tyr) were 5 preferentially released and a typical cocoa aroma was detected after roasting in the presence of reducing sugars (Voigt et al., Food Chem. 50 (1994), 177-184). In contrary to that, the predominant amino acids released from the albumin-derived peptides were aspartic acid, glutamic acid and asparagine. Furthermore, no cocoa aroma was detected with the albumin fraction. It was therefore concluded that cocoa-specific aroma precursors are preferentially 10 derived from the vicilin-like globulin of cacao bean. Consequently, the mixture of hydrophobic free amino acids and remaining oligopeptides required for the generation of the typical cocoa flavour components seems to be determined by the particular chemical structure of the cacao vicilin-class globulins.

15 These globulins isolated from cacao beans were also found to be efficiently degraded by pepsin (an aspartic endoproteinase) and chymotrypsin (a serine endoproteinase). Products derived from cacao globulins by successive proteolytic digestion with pepsin and carboxypeptidase A revealed a typical, but less pronounced cocoa aroma upon roasting. No cocoa aroma precursors were generated by degradation of globulins with chymotrypsin and 20 carboxypeptidase A (Voigt et al., Food chem, 51 (1994), 7-14). Therefore, the specific mixture of oligopeptides and hydrophobic free amino acids required for the formation of the typical cocoa aroma is not only determined by the structure of the protein substrate but also dependent on the specificity of the cacao enzyme cleaving the protein.

25 In view of the above data a hypothetical model for the generation of the said mixture of peptides and amino acids, i.e. the cocoa flavour precursors, during fermentation had been devised (Fig.1), wherein in a first step peptides having a hydrophobic amino acid at their end, are formed from storage proteins, which peptides are subsequently further degraded. For splitting off hydrophobic amino acids from peptides formed in a preceding step the above 30 carboxypeptidase activity seems to be involved. Yet, for the stage of producing the said peptides having C-terminal hydrophobic amino acids, the only known enzymatic activity

which might be considered in this respect is an aspartic endoproteinase activity related to that mentioned above. It is also possible that the activity mentioned above is the result of different enzyme activities which are still unknown.

5 Though some aspects of cocoa flavour production have been elucidated there is still a need in the art to fully understand the processes going on, so that the manufacture of cocoa flavour may eventually be optimized.

An object of the present invention therefore resides in providing means to improve the

10 formation of cocoa flavour during processing and manufacturing.

The above object has been solved by providing two novel aspartic endoproteinases derived from *Th. cacao* as identified by SEQ ID No 1 and SEQ ID No 2 or variants thereof obtained by substituting, deleting or adding one or more amino acids such, that the enzymatic activity

15 thereof is essentially retained. The aspartic endoproteinases described here (termed TcAP1 and TcAP2 in the following) shall be capable to cleave the vicilin-class globulins isolated from cacao beans so that a successive degradation of the peptides by means of carboxypeptidase will result in a mixture of peptides and amino acids that yields a cocoa flavour upon a reaction with reducing sugars, i.e. upon roasting.

20

According to another embodiment the present invention provides DNA sequences coding for the respective endoproteinases. The DNA sequences may be derived according to the genetic code from the amino acid sequences as identified under SEQ ID Nos. 1 and 2 considering the wobble hypothesis, optionally taking account codon preferences of specific hosts, in which

25 the DNA sequences shall be expressed. The skilled person may well devise appropriate DNA sequences based on the polypeptide sequences given and his own technical knowledge and understanding. According to a preferred embodiment the DNA sequences are as identified under SEQ ID No 3 (TcAP1) and SEQ ID No 4 (TcAP2), which DNA sequence may be varied by replacing, deleting or adding one or more nucleotides such, that the endoproteinases essentially retain their enzymatic activity.

The DNA sequences may be used for recombinantly preparing the aspartic endoproteinases of the present invention. To this end the DNA sequences are incorporated into a suitable expression vector, such as a plasmid or a viral vector, which comprise the common sequences, such as a promotor, a polylinker for alleviating the cloning of the DNA sequences 5 therein, leader sequences, to direct the polypeptide produced out of the cell. The vectors will be selected based on the requirements of the system used, e.g. for an expression in *E. coli* the vectors pGEMEX, pUC-derivates, pGEX-2T, pET-derivates, pQE8 may be envisaged, which are widespread in use and are commercially available. As an example aspartic endoproteinases could be expressed into medium or on surface of lactic acid bacteria used in lactic 10 products such as milk or yogurt.

For expressing the endoproteinases in e.g. yeast the vectors pNFF296, pY100, pPIC9K, pPICz and Ycpad1 may be utilized and for expression in animal cells the vectors pKCR, pEFBOS, cDM8 und pCEV4 as well as pSS-derivates (Kay R. et al., Science 236 (1987), 15 1299-1302) may be used. Moreover, for expressing the endoproteinases in plant cells, especially in cacao, the vector pAL76 or pBin19-derivates may be used and for insect cells e.g. the vector pAcSGNT-A.

The aspartic endoproteinases may be expressed in a prokaryotic or eukaryotic cell as 20 mentioned above. It will be appreciated that the skilled person will be able to select, based on the need and his own technical skill, an appropriate expression system to achieve the desired goal. In case the endoproteinase shall simply be added to a protein mixture, such as isolated cacao vicilin-class globulins, the recombinant enzyme may be produced in a bacterial system such as *E. coli* or in yeast and applied on the protein material.

25 Yet, in view of the implication to increase the enzymatic activity in cacao itself a transgenic plant cell may be envisaged, wherein one or more copies of the endoproteinases, optionally coupled with a suitable and controllable promotor, have been incorporated into the genome of the plant cell. The introduction of the DNA sequence(s) may be achieved by e.g. 30 homologous recombination of DNA stretches harboring one or more copies of the DNA sequences coding for the endoproteinases of the present invention into embryogenic calli

prepared beforehand. Since plant cells are totipotent a new transgenic cacao tree may be produced in this way the beans of which will exhibit more rapid degradation of the vicilin-class globulins when subjected to conditions of fermentation.

5 In consequence a transgenic plant, harboring one or more additional copy(ies) of a DNA sequence coding for the endoproteinases of the present invention is well within the scope of the present invention.

The present endoproteinases may also be used for the manufacture of cocoa flavour by
10 treating a suitable starting material (cacao bean, liquor or crumb), preferably vicilin-class globulins, with said endoproteinases of the present invention and concurrently or afterwards treating the material with carboxypeptidase to obtain a mixture of peptides and amino acids appropriate to act as cocoa flavour precursors. This mixture may then be subjected to "a roasting step", i.e. may be subjected to a reaction with reducing sugars to eventually obtain
15 cocoa flavour.

Since some of the enzymes involved in the generation of cocoa flavour are now at hand
cocoa flavour may be produced artificially without having to rely on the common process of
fermenting and roasting cacao beans. The present invention therefore also provides a method
20 for generating cocoa flavor which comprises the step of subjecting a material suitable to yield
cocoa flavour precursors, such as the known vicilin-class globulins, to an enzymatic
degradation involving the use of the aspartic endoproteinases of the present invention.

In particular, the present aspartic endoproteinases may be overexpressed in protein bodies of
25 plant cells, especially seed cells, and then hydrolysis of the cellular protein material may be
effected by treating such plant cells with an acidic solution.

The present endoproteinases may also be used for hydrolyzing proteins by contacting a
material of choice, such as the protein in isolated form or material containing the protein,
30 such as e.g. food material, with an endoproteinase of the present invention and effecting
hydrolysis to a desired degree. Examples for materials are dairy substances (whey protein,

and casein), wheat gluten, corn gluten, meat, egg protein and other protein containing vegetable substances not mentioned above such as proteins from oil seeds, including soybean protein and defatted soy protein.

5 In the figures,

Fig. 1 shows the theoretical production process of cocoa specific flavour precursors;

10 Fig. 2 shows a schematic representation of plant aspartic prepropeptides;

15 Fig. 3 schematically shows the cloning strategy for the isolation of the endoproteinase *TcAPI* cDNA;

Fig. 4 schematically shows the cloning strategy for the isolation of the endoproteinase *TcAP2* cDNA;

15 Fig. 5 shows a comparison between the different polypeptides obtained;

Fig. 6 shows a hydrophilicity Kyte Doolittle plot for both endoproteinases obtained.

20 Fig. 7 shows the expression of *TcAPIa* and *TcAP2* in cacao beans of three different cacao clones;

25 Fig. 8 shows the results of a Northern blot analysis of *TcAPIa* and *TcAP2* expression in cacao beans from clone CCN51 at different maturation stages;

Fig. 9 shows the results of a Northern blot analysis of *TcAPIa* and *TcAP2* expression in cacao bean produced by cacao clone CCN51 at different germination stages;

30 Fig. 10 shows the results of a hydrolysis experiment of bovine haemoglobin by recombinant *TcAP2* protein in yeast culture medium and comparison with control strain pNFF296;

Fig. 11 shows the results of experiments determining the pH dependence of haemoglobin hydrolysis by recombinant TcAP2;

5 Fig. 12 shows the effect of different inhibitors on the hydrolysis of bovine haemoglobin by recombinant TcAP2;

10 Fig. 13 shows the analysis of most active pool (fractions 57-64) from the Sephadryl S-200 HiPrep 16/60 size exclusion column on a 10-20% Gradient SDS-PAGE Gel (Coomassie 15 stained). In lanes 1-3, 12, 24, and 40.8 µg protein was loaded in each lane respectively. Complex denotes a putative covalent complex between AP and trypsin inhibitor fragments; TcAP2 denotes the 30.5 kDa polypeptide; 27.9 denotes the 27.9 kDa putative endochitinase; TI, trypsin inhibitor. The molecular weights of the markers are noted on the right;

15 Fig. 14 shows SDS-PAGE gel analysis of the reaction products after a Q Sepharose Fast Flow purified aspartic endoproteinase preparation was incubated in acid conditions for 1 minute and 7 hours. AP denotes the 30.5 kDa polypeptide; 27.9 denotes the 27.9 kDa putative endochitinase; TI, trypsin inhibitor. M, molecular weight markers (Precision, Biorad);

20 Fig. 15 shows denaturing size exclusion chromatography of the reaction products after a Q Sepharose Fast Flow purified aspartic endoproteinase was incubated in acid conditions for 1 minute and 7 hours respectively. The molecular weight size markers are: 1, ribonuclease A 13.7 kDa; 2, aprotinin 6.5 kDa; 3, substance P 1,347 Da; 4, N-benzoyl-gly-phe (hippuryl-phe) 326 Da; 5, phe 165 Da;

During the studies leading to the present invention two aspartic endoproteinases have been found which seem to participate in the enzymatic degradation of vicilin-class globulins in cocoa beans under the conditions of fermentation.

30 Aspartic endoproteinases as such are a widely distributed class of proteases in animals,

microbes, viruses and plants. All aspartic endoproteinases contain two aspartic residues at the active site and are active at acidic pH. In most of the aspartic endoproteinases, the catalytic aspartic residues are contained in a common Asp-Thr-Gly motif present in both lobes of the enzyme, with plant aspartic endoproteinases containing Asp-Ser-Gly at one of the sites.

5

Many aspartic proteinases have been detected or purified in monocots and dicots, which are either heterodimeric or monomeric. The sequences of corresponding genes predict that the active heterodimeric enzymes are derived from the processing of a single proprotein.

10 Though the genes and predicted proproteins for both monomeric and dimeric plant aspartic endoproteinases are quite similar they differ from mammalian and microbial counterparts by the presence of a 100 amino acids insert (a so called plant specific insert: PSI) which is absent in mammalian and microbial aspartic proteinases. This insert divides the protein in two regions: an amino-terminal and a carboxy-terminal region which show a relatively high
15 similarity to each other and to mammalian and microbial enzymes. The amino-terminal region contains the two active sites Asp-Thr-Gly (DTG) and Asp-Ser-Gly (DSG) (Fig 2). Although the positions of six cysteine residues are conserved, the PSI from different species are less homologous with each other than are the amino- and carboxy-terminal regions.

20 In view of this knowledge the conserved region has been utilized to obtain the nucleotide and amino acid sequence of aspartic endoproteinase (TcAPI) from cacao bean (clone ICS 95) as follows:

A 1 kb internal fragment of the aspartic proteinase from cacao bean was amplified by RT-
25 PCR using degenerate oligonucleotides that had been chosen according to an alignment of known aspartic endoproteinase sequences and a selection of conserved regions. Based on the sequence of this fragment, primers were designed to amplify 5'- and 3'-end. Afterwards, a full-length cDNA (*TcAPIb*) was obtained by ligation of the 3' and 5' fragment using the *Bam*H I restriction site and another one (*TcAPIa*) was amplified using primers specific to
30 both extremities (Fig. 3).

TcAP1a and *TcAP1b* nucleotide sequences differ only by 6 base pairs. Some of these differences are also found in the partial 1 kb fragment. Three of the differences lead to amino acid changes in the encoded protein (Table 1). The molecular weight and the pI of the protein are not changed.

5

Table 1. Differences observed in the nucleotide sequences from the different cDNA fragments obtained by PCR and their impact on the protein sequence.

Position	1 kb fragment	TcAP1a	TcAP1b	Altered residue
318		T	A	L----M
431	C	C	T	No change
636	G	G	A	A----T
764	T	C	T	No change
1189	C	T	C	V----A
1376	C	C	T	No change

10

These differences may be explained by mistakes performed by polymerase enzymes during the PCR reactions. Another explanation could be that *TcAPIa* and *TcAPIb* are two different alleles from the same gene that we will name *TcAPI*. Furthermore, the 5'- and 3'-untranslated regions from *TcAPIa* and *TcAPIb* are identical. This argues rather for the presence of two alleles than for two different genes.

15

The cDNA sequences from *TcAPIa* (SEQ ID No 3) isolated from cacao bean (clone ICS95) is 1784 bp long. A putative initiation start codon was assigned by comparison with other plant aspartic proteinase sequences. It is located 63 bp from the 5' end. The open reading frame is broken by a stop codon (TAA) at position 1605, followed by a putative polyadenylation signal (TATAAAA) at position 1625.

TcAP1a encodes a 514 amino acid protein with a predicted molecular weight of 56 kDa and a pI of 5.05. The protein shows a high similarity with plant aspartic endoproteinasies.

25

Considering entire sequences, percent identity ranged between 59% observed with rice aspartic endoproteinase (Oryzasin A) and 87% with partial cotton sequence. A

hydrophobicity analysis (Fig. 6A) reveals that *TcAP1a* encodes a hydrophilic protein with a very hydrophobic N-terminal end, indicating the presence of a signal peptide. Two catalytic triads (DTG and DSG) are also present.

5 The nucleotide and amino acid sequence of aspartic endoproteinase (*TcAP2*) from cacao bean (clone CCN51) was obtained as follows:

A 1 kb internal fragment of the aspartic endoproteinase from cacao bean was amplified by RT-PCR using degenerate oligonucleotides selected as above. Based on the sequence of this 10 fragment, primers were designed to amplify 5'- and 3'-end. Afterwards, a full-length cDNA (*TcAP2*) was amplified using primers specific to both extremities (Fig. 4).

The cDNA sequence from *TcAP2* (SEQ ID No 4) isolated from cacao bean (clone CCN51) is 1828 bp long. An initiation start codon is located 62 bp from the 5' end. The open reading

15 frame is broken by a stop codon (TAA) at position 1606, followed by a putative polyadenylation signal (TATAAA) at position 1669.

TcAP2 encodes a 514 amino acid protein with a predicted molecular weight of 56 kDa and a pI of 5.31. The protein shows a high similarity with plant aspartic endoproteinases.

20 Considering entire sequences, percent identity ranged between 57% observed with rice aspartic endoproteinase (Oryzasin A) and 77% with partial cotton sequence. A hydrophobicity analysis (Fig 6B) reveals that *TcAP2* encodes a hydrophilic protein with a very hydrophobic N-terminal end, indicating the presence of a signal peptide. Two catalytic triads (DTG and DSG) are also present.

25

The following examples illustrate the invention without limiting it to the same.

Cacao (*Theobroma cacao L.*) beans from ripe pods of clones ICS 95, CCN51 and EET95 were provided by Nestlé ex-R&D Center Quito (Ecuador). The beans were taken from the

30 pods immediately after arrival at the laboratory (4-5 days after harvesting). The pulp and the seed coat were eliminated and the cotyledons were frozen in liquid nitrogen and stored at -

80°C until use.

Example 1

Preparation of mRNA

5

Two beans were grounded in liquid nitrogen to a fine powder and extraction was directly performed with a lysis buffer containing 100 mM Tris HCl pH8, 1% SDS and 0.1M β -mercaptoethanol. RNA was extracted with one volume phenol/chloroform/isoamylalcohol (25/24/1) and centrifuged at 8000 rpm for 10 min at 4°C. The aqueous phase was washed three times with chloroform/isoamylalcohol (24/1). RNA was precipitated with 0.3M sodium acetate pH 5.2 in two volumes of ethanol. The RNA pellet obtained after centrifugation was resuspended in 100 mM Tris HCl pH 8 and a second precipitation with 2M lithium chloride was performed. The RNA pellet was washed with 70% ethanol and resuspended in DEPC treated water.

15

Example 2

Cloning of aspartic proteinase cDNAs

20 A search for aspartic proteinase sequences in the GenBank database led to the identification of several plant sequences. A multiple alignment of these sequences revealed the presence of conserved regions, which have been used to design two degenerate oligonucleotides:

25 A sense primer, pAP0 (5'-GAYACNGGNAGYTCYAAYYTVTGG) has been synthesised according to the sequence Asp-Thr-Gly-Ser-Ser-Asn-Leu-Trp, which contains an active site (Asp-Thr-Gly) of the protein.

An antisense primer pAP4r (5'-CCATMAANACRTCNCCMARRATCC) has been synthesised according to the sequence Trp-Ile-Leu-Gly-Asp-Val-Phe, located in the C-terminal part of the protein.

30

Total RNA as prepared in example 1 was used to synthesize first strand cDNA with the

SMART PCR cDNA Synthesis Kit (Clontech, USA). Synthesis has been performed exactly as described in the kit instructions using 1 μ g of total RNA and the SuperscriptTM II MMLV reverse transcriptase (Gibco BRL, USA). After synthesis, cDNA was used directly for PCR or kept at -20°C.

5

Specific cDNA amplification was performed with 2 μ l first strand cDNA in 50 μ l buffer containing 10mM Tris-HCl pH 8.8, 50mM KCl, 1.5 mM MgCl₂, 0.001 % (w/v) gelatin, 0.25 mM dNTP's, 30 pmoles of pAP0 and pAP4r primers and 5 units of Taq DNA polymerase (Stratagene, USA). Amplification was performed in a Bio-med thermocycler 60 (B. Braun).

10 A first denaturation step (94°C, 2 min) was followed by 30 cycles of denaturation (94°C, 1 min), primer annealing (40°C, 1.5 min) and extension (72°C, 2 min). The extension time was increased by 3 sec at each cycle. Amplification was ended by a final extension step (72°C, 10 min). The amplified fragment was cloned in pGEM[®]-T Easy vector and sequenced.

15 *TcAPI* and *TcAP2* full-length cDNAs were cloned using Rapid Amplification cDNA Ends PCR (RACE PCR). For *TcAPI*, the MarathonTM cDNA Amplification Kit (Clontech, USA) was used. Poly A+ RNA purified from total RNA (150 μ g) with the Oligotex mRNA kit (QIAGEN, Germany) were used for the synthesis of double strand cDNA and a Marathon cDNA adaptor was ligated at both ends of the cDNA. These two steps have been performed 20 according to the instructions of the MarathonTM cDNA Amplification Kit. For *TcAP2*, single strand cDNA has been synthesised from total RNA according to the SMARTTM RACE cDNA Amplification Kit (Clontech, USA).

25 RACE PCR was performed with 5 μ l Marathon adaptor-ligated double strand cDNA or 2.5 μ l SMART single strand cDNA in 50 μ l buffer containing 40 mM Tricine-KOH pH 9.2, 15 mM KOAc, 3.5 mM Mg(OAc)₂, 3.75 μ g/ml BSA, 0.005% Tween-20, 0.005% Nonidet-P40, 0.2 mM dNTP's, 0.2 μ M of each primer and 1 μ l Advantage 2 Polymerase mix (Clontech, USA). Amplification was performed via touchdown PCR, in a Bio-med thermocycler 60 (B. Braun).

30

A first denaturation step (94°C, 1 min) was followed by:

- 5 cycles including denaturation at 94°C for 30 sec and annealing/extension at 72°C for 7 min
- 5 cycles including denaturation at 94°C for 30 sec and annealing/extension at 70°C for 7 min
- 5 - 25 cycles including denaturation at 94°C for 20 sec and annealing/extension at 68°C for 7 min

10 For *TcAP1*, two specific primers were paired with the AP1 primer, specific to the Marathon cDNA Adaptor provided in the Marathon kit:

ICS5 for 5'RACE PCR reaction (5'GCAGGCCACCAGCACAAAGTCCAG)

ICS3 3'RACE PCR reaction (5'CGGTTGGAAATGCTGTGCCTGTGTGG)

15 For *TcAP2*, two specific primers were paired with the UPM (Universal Primer Mix) primer that recognises the SMART sequence:

CCN5 for the 5'RACE PCR reaction (5'ATGTGTGCTTGCCCTTGTAGTGG)

CCN3 for the 3'RACE PCR reaction (5'CCGCAATGTAGATGAAGAAGCAGGTGG)

20 The amplified fragments were cloned in pGEM®-T Easy vector and sequenced. The sequence information obtained after the sequencing of RACE fragments was used to design new oligonucleotides in order to amplify the full length fragments:

25	TcAP1	TcAP1, sense primer (5'TCTGCTCAGCTTTCTTGTGCG)
		TcAP1r, reverse primer (5'GGATCACATGAAATTCTAAACAAAGTGC).
	TcAP2	TcAP2, sense primer (5'CTAACATCGACTCACTATAGG)
		TcAP2r, reverse primer (5'ATCTGTGACTGTTGATAAAAAGC)

30 PCR reaction was performed exactly as for the amplification of 5'- and 3'-RACE fragments with one denaturation step (95°C, 1 min) followed by 35 cycles of denaturation (94°C, 30

scc), primer annealing (63°C, 1 min) and extension (72°C, 2 min). The extension time was increased by 3 sec at each cycle. Amplification was ended by a final extension step (72°C, 10 min). The amplified fragment TcAP1 and TcAP2 were cloned in pGEM®-T Easy or pGEM®-T vectors respectively and sequenced.

5

Furthermore, a cloning strategy was also used to obtain the full-length TcAP1 cDNA. 5'- and 3'-RACE fragments overlap for 200 base pairs. In this overlapping region an unique restriction site BamH I is present. Both fragments have been isolated using *Bam*H I and *Eco*R I (present in the plasmid) and subcloned directly in pBS+ (Stratagene, USA) using the same restriction enzymes.

Example 3

Sequencing and analysis of DNA sequences

15 cDNA sequencing has been performed according to standard techniques (Maniatis, A Laboratory Manual, Cold Spring Harbor, 1992). Sequence analysis and comparison were done using DNASTar programme. The sequences are shown under SEQ ID Nos 1 and 2.

Example 4

20 Expression of *TcAP1a* and *TcAP2* in cacao plants

For the Northern blot total RNA was separated on 1.5 % agarose gel containing 6% formaldehyde in 20mM MOPS, 5mM NaOAC, 1mM EDTA pH 7. After electrophoresis, RNA was blotted onto nylon membranes (Appligene) and hybridized with ³²P-labeled 25 *TcAP1a* or *TcAP2* probe at 65°C in 250mM Na-phosphate buffer pH 7.2, 6.6% SDS, 1 mM EDTA and 1% BSA. Membranes were washed three times at 65°C for 30 min in 2XSSC, 0.1%SDS; in 1XSSC, 0.1% SDS and finally in 0.5XSSC, 0.1%SDS.

30 *TcAP1a* probe was amplified by PCR using TcAP1 and TcAP1r primers and *TcAP2* probe with the following primers:

TcAP2b: a sense primer (5'-CTATAGGGCAAGCAGTGGTAACAAAC)
TcAP2br: an antisense primer (5'-TGACCTAAAGGCAAATCCTAGTTTC)

PCR reaction was performed with 1 μ l template cDNA in 50 μ l buffer containing: 40 mM 5 Tricine-KOH pH 8.7, 15 mM KOAc, 3.5 mM Mg(OAc)₂, 3.75 μ g/ml BSA, 0.005% Tween-20, 0.005% Noninet-P40, 0.2 mM dNTP's, 0.2 μ M of each primer and 1 μ l 50X Advantage 2 20 polymerase Mix (Clontech, USA). Amplification was performed in a Bio-med thermocycler 60 (B. Braun). A first denaturation step (94°C, 1 min) was followed by 30 cycles of denaturation (94°C, 30 sec), primer annealing (63°C, 1.5 min) and extension (72°C, 2 min). 10 The extension time was increased by 3 sec at each cycle. Amplification was ended by a final extension step (72°C, 10 min).

Both fragments were purified with Strataprep PCR purification kit (Stratagene, USA) and 15 labelled by the random priming procedure (*rediprime*TM II, Amersham Pharmacia Biotech).

15 Northern blot analysis with RNA purified from mature cacao beans produced by different trees, CCN51, EET95 and ICS95 reveals that *TcAP1a* and *TcAP2* are both expressed in beans produced by the three different trees (Fig. 7A). However, *TcAP2* is much more strongly expressed than *TcAP1a* indicating that it might be the major aspartic endoproteinase 20 in cacao beans. RT-PCR experiments (Fig. 7B) are in agreement with these results. Confirmation of the idea that *TcAP2* is the major aspartic endoproteinase activity in the bean is provided by the N-terminal sequencing of a purified native protein, which has the same sequence than *TcAP2*. Finally, the RT-PCR results presented in figure 7B also clearly show that both genes are expressed in leaves.

25 Similar experiments performed with RNA purified from cacao beans at different stages of maturation (Fig. 8) confirm that *TcAP1* is less expressed than *TcAP2* in developing and mature beans. *TcAP1* and *TcAP2* expression increase slightly during maturation and decrease in mature beans. *TcAP2* is mainly expressed in early bean developmental stages suggesting 30 that the synthesis of new aspartic endoproteinase falls as the bean matures.

During germination, the expression of *TcAP2* is relatively stable in contrary to that of *TcAP1*, which increases after a few days of germination with a maximum at days 4 and 7. A strong expression is also detected at 49 days after imbibition (Fig. 9).

5 **Example 5**

cDNA expression in yeast heterologous system

The coding sequences of *TcAP1a* and *TcAP2* were overexpressed in the yeast heterologous system *Yarrowia lipolytica*.

10

TcAP1a and *TcAP2* were overexpressed under the control of a synthetic XPR2-derived promoter hp4d present on the *Yarrowia lipolytica* expression/secretion plasmid pNFF296. For both cDNA, in order to excrete the recombinant protein in the culture medium the signal sequence (first 24 amino acids, predicted as according to Nielsen et al., Protein Engineering 15 10 (1997), 1-6 was replaced by a lipase signal sequence present on the *Yarrowia lipolytica* expression/secretion plasmid pNFF296.

TcAP1a cloned in pGEM-T Easy was used as template for the amplification of the cDNA sequence coding for a mature protein without a putative signal sequence.

20

Two primers were used for the amplification of *TcAP1a*:

Primer C089

(5'-CCGGCCTTCGGCCGCCAAGCGAATATCCAATGAGAGATTGGTCAG)

25

primes at the 5' end of the predicted mature *TcAP1a* cDNA and introduces a SfiI site allowing cloning in frame to a hybrid XPR2-lipase signal sequence present on the *Yarrowia lipolytica* expression/secretion plasmid pNFF296

Primer C090

30 (5'-CCGGCCCACGTGGCCTAGTGGTGGTGTGCAGCCTGGCAAATCCAAC)
primes at the 3' end of the mature *TcAP1a* cDNA and introduces in-frame a 3xHIS sequence

just before the stop codon and the SfiI cloning site in front of the lipase terminator of pNFF296.

5 TcAP2 cDNA cloned in pGEM-T was used as template for the amplification of the sequence coding for the mature protein without a putative signal sequence.

Two primers were used for the amplification of TcAP2:

Primer C091

10 (5'-CCGGCCTCTCGGCCAAGCGAGTATCCAATGATGGGCTGGTAG) primes at the 5' end of the predicted mature TcAP2 cDNA and introduces a SfiI site allowing cloning in frame to a hybrid XPR2-lipase signal sequence present on the *Yarrowia lipolytica* expression/secretion plasmid pNFF296.

15 Primer C092

(5'-CCGGCCCACGTGGCCTAGTGGTGGTGTGCCGCCTCGCGAAGCCGAC) primes at the 3' end of the mature TcAP2 cDNA and introduces in-frame a 3xHIS sequence just before the stop codon and the SfiI cloning site in front of the lipase terminator of pNFF296.

20

Amplification was performed with 1 μ l of template cDNA (20 ng) in 10 mM KCl, 6 mM $(\text{NH}_4)_2\text{SO}_4$, 20 mM Tris-HCl, pH 8.0, 0.1% Triton X-100, 2 mM MgCl₂, 0.2 mM of each dNTP, 10 μ g ml⁻¹ BSA, 0.25 μ M of each primers and 3 units of Pfu DNA polymerase (Stratagene, USA). PCR was performed in a Stratagene RoboCycler (Stratagene, USA). A 25 first cycle (95°C-5 min, 50°C-1 min, 72°C-3 min) was followed by 30 cycles (95°C-1 min, 50°C-1 min, 72°C-3 min) and a final cycle (95°C-1 min, 50°C-1 min, 72°C-10 min). The PCR products were purified using the Qiaquick PCR purification Kit (Qiagen INC., USA), digested with SfiI, and subsequently ligated into vector pNFF296 previously digested with SfiI. This ligation was used to transform *E. coli* BZ234 (Biozentrum, University of Basel, 30 Switzerland). Constructs were selected on LB plates supplemented with 50 μ g ml⁻¹ kanamycin, analyzed by mini plasmid-preparations plus restriction enzyme digestion and

finally by DNA sequence analysis. The resulting plasmids containing TcAP1a or TcAP2 were called pCY329 and pCY330, respectively.

The *Yarrowia lipolytica* host strain YLP3 was derived from strain polf (MatA ura3-302 5 leu2-270 xpr2-322 axp-2 SUC2) by transforming said strain to leucine prototrophy with a 5.1 kb SalI fragment carrying the *Yarrowia lipolytica* wild-type LEU2 gene (J.-M. Nicaud, pers. comm.) and selecting for LEU2 convertants. The *Yarrowia lipolytica* host strain was streaked on a YPD agar plate (1% Difco Bacto Yeast Extract, 2% Difco Bacto Peptone, 2% Glucose, 10 2% Difco Bacto Agar) and grown overnight at 28°C. 4 ml of liquid YPD pH 4.0 (1% Difco Bacto Yeast Extract, 1% Difco Bacto Peptone, 1% Glucose, 50 mM Citrate buffer at pH 4.0) were inoculated with freshly grown cells of the YPD plate and grown in a tube on a rotary shaker (200 rpm, 28°C, 8-9 hrs). Of this preculture an adequate amount was used to inoculate 20 ml YPD pH 4.0 in a 250 ml Erlenmeyer flask without baffles. This culture was shaken in a 15 rotary shaker at 200 rpm at 28°C (over night) until a cell titration of 10^8 ml⁻¹ has been reached. The cells were centrifuged for 5 min at 3000 g, washed with 10 ml of sterile water and re-centrifuged. The cellular pellet was suspended in 40 ml 0.1 M lithium acetate pH 6.0 (adjusted with 10% acetic acid) and shaken in a 250 ml Erlenmeyer at 140 rpm at 28°C for 60 minutes. The cells were again centrifuged for 5 min at 3000 g. The cellular pellet was suspended in 2 ml lithium acetate pH 6.0 and the competent cells were kept on ice until 20 transformation.

One hundred microliters of competent cells were mixed with 5-20 µl plasmid linearized with 25 NotI and 50 µg carrier DNA (herring sperm DNA sonicated to 100-600 bp, Promega, USA) in a 2 ml tube and incubated for 15 minutes at 28°C. 700 µl 40% PEG4000, 0.1 M lithium acetate pH 6.0 were added and the tubes heavily agitated at 240 rpm on a rotary shaker at 28°C for 60 minutes. A volume of 1.2 ml of 0.1 M lithium acetate pH 6.0 was added and mixed. 250 µl were plated on selective agar plates (0.17% Difco Bacto Yeast Nitrogen Base w/o amino acid and ammonium sulfate, 1% glucose, 0.006% L-leucine, 0.1% sodium glutamate, 0.1% Difco Bacto Casamino Acids, 2% agar). The expression plasmid pNFF296 30 carries a defective URA3 allele allowing for the selection of multiple integration of the

expression secretion cassette in the YLP3 host strain.

Transformants (Ura^+) were re-isolated on selective medium (0.17% Difco Bacto Yeast Nitrogen Base w/o amino acid and ammonium sulfate, 1% glucose, 0.006% L-leucine, 0.1% sodium glutamate, 0.1% Difco Bacto Casamino Acids, 2% agar). A series of clones was grown in shake-flasks to check for expression and secretion of aspartic proteinase into the culture medium.

Small patches of cells were streaked on YPD agar plates and grown overnight at 28°C. The 10 thin layers of grown cells were used to inoculate 50 ml DMI medium in 500 ml Erlenmeyers with 4 lateral baffles. DMI medium contains per liter: KH_2PO_4 , 10 g; $MgSO_4 \cdot 7H_2O$, 2.5 g; glucose, 20 g; Trace elements solution, 5.1 ml; Vitamins solution, 17 ml; urea, 3 g. Urea was dissolved in 15 ml water and sterile filtered. The initial pH of the medium was adjusted to 5.0. The cultures were shaken at 140 rpm on a rotary shaker at 28°C for three days. Aliquots 15 of the cultures were centrifuged at maximum speed (3000 g) for 15 min. and the supernatant used for the determination of the aspartic endoproteinase activity.

Aspartic endoproteinase activity was assayed at 42°C in a 900 μ l reaction medium containing 0.2M sodium citrate buffer pH3.0, 10 mg/ml bovine haemoglobin and 150 μ l yeast culture 20 supernatant. To stop the reaction aliquots (80 μ l) were added to an equal volume of TCA 8% and the precipitated protein removed by centrifugation at 13000 g. 20 μ l supernatant were mixed to 250 μ l O-phthaldialdehyde (OPA) reagent (50 mM sodium tetraborate, 1 % SDS, 5.96 mM OPA (dissolved in 1 ml methanol) and 1.43 mM β -mercaptoethanol. Activity was then determined measuring OD at 340 nm and expressed in pmole leucine produced per mg 25 protein. For this, we use the following linear equation ($OD_{340nm} = 0.0156 \text{ pmoles} + 0.0088$), which was determined using a standard curve with L-leucine (0 to 80 pmoles). Protein concentration was determined by Bradford assay (Biorad).

A strong activity could be detected in 12 independent clones transformed with the pCY330 30 construct (TcAP2). Further characterization of the TcAP2 recombinant protein was done using one clone named pCY330-33. Comparison of activity measurement with supernatant

from pCY330-33 and pNFF296 (control) clearly shows that no activity is detected in the control (1.44 ± 0.52 pmoles L-leucine/min/mg protein) and that hydrolysis of bovine haemoglobin occurs in presence of supernatant from pCY330-33 (25.8 ± 1.45 pmoles L-leucine/min/mg protein) (Fig. 10). This activity demonstrates clearly that active recombinant

5 TcAP2 protein is produced by pCY330-33.

The recombinant TcAP2 endoproteinase detected in pCY330-33 hydrolyses bovine haemoglobin with an optimum at pH 3 (Fig. 11). Only slight activity could be detected for pH higher than 5.

10

The endoprotease activity detected in the medium of pCY330-33 (TcAP2) is completely inhibited by 2 μ M pepstatin, a specific inhibitor for aspartic endoproteinase. The pepstatin insensitive activity (1.91 ± 1.26 pmoles L-leucine/min/mg protein, 6.65%) is in the same range as that one measured for the control strain (2.26 ± 1.26 pmoles L-leucine/min/mg protein, 7.8%). Other inhibitors such as 1.10 phenanthroline (metallo proteases), DCI (serine proteases) and E64 (cysteine proteases) have no effect on TcAP2 activity (Fig. 12).

15

The data presented here clearly show that the culture medium in which yeast pCY330-33 was grown contained a protein able to hydrolyse bovine haemoglobin. Maximum activity at acidic pH and inhibition by pepstatin are two specific biochemical features for aspartic proteinases.

Example 6

Native protein purification

25

Approximately 25 g of the frozen EET 95 cacao beans were ground to a fine powder using liquid nitrogen and extracted with cold acetone/water/5mM sodium ascorbate (80/20/5) according to a modified procedure of Hansen et al., J. Sci. Food Agric. 77 (1998), 273-281, to remove the majority of the fat and phenolic compounds. This procedure resulted in approximately 11.3 g of a

30 fine acetone powder.

Acetone powder (5g) was extracted twice with 500 ml of buffer A (10 mM sodium phosphate pH 7.8, 2 mM EDTA, 10 mM sodium acetate) for 1 hour at 4°C. After centrifugation (7840g, 25 min, 4°C) the combined supernatants were made sequentially to 30% and 60% ammonium sulphate. All ammonium sulphate fractions were assayed for activity and the 60% ammonium sulphate precipitate was found to have the highest level of endoproteinase activity and was dialysed against buffer B (50 mM sodium phosphate pH 7.8, 1 mM EDTA).

Using an Akta Purifier (Pharmacia), 2 x 10 ml of dialysed 60% ammonium sulfate precipitate were loaded on a HiLoad 26/10 Q Sepharose Fast Flow column (Pharmacia) at 8-10°C. After 10 loading, the column was washed with 5 column volumes of 20 mM Tris-HCl pH 8, then eluted with a linear gradient of 10 column volumes of the same buffer supplemented with 1 M NaCl. The flow rate of the column was 10 ml/min and 5 ml fractions were collected.

Fractions from the Q Sepharose Fast Flow column were assayed for aspartic endoproteinase 15 activity and fractions showing the highest level of activity (#65-80) were pooled. The pooled fractions (75 ml) were concentrated to 2.2 ml using "Ultrafree Biomax" 4 ml filters (5 kDa Mw cut off), and loaded onto a Sephadryl S-200 HiPrep 16/60 size exclusion column (Pharmacia) equilibrated with 10mM Tris-HCl pH 8 and 500 mM NaCl at a flow rate of 0.5 ml/min. 1 ml 20 fractions were collected and assayed for aspartic endoproteinase activity. The most active fractions were concentrated into three pools (#53-56, #57-64, #65-68) using "Ultrafree Biomax" filter. Protein concentration was determined with the micro BCA protein assay kit (Pierce, Inc) using BSA as a standard.

The most active pool (#57-64) with a specific activity of 1054 units/mg protein (1unit=100 ng 25 leucine equivalent produced/min) has been subjected to SDS-PAGE. This gel (Fig. 13) shows that this fraction contains several polypeptides. N-terminal sequencing of the major bands revealed that only the 30.5 kDa band (DSEETDIVAL) corresponded exactly to the sequence of the cacao TcAP2 protein of the present invention. The other main polypeptides in the preparation were found to be putative protein body proteins. The 27.9 kDa polypeptide N-terminal sequence 30 (TVIYSTYWGQNGFEGT) showed the strongest homology (76.9%) with a *Glycine max* acid chitinase III-A (accession AB007127). Thus, it is likely that the 27.9 kDa protein is an acid chitinase. The N-terminal sequence obtained for the 20.2 kDa polypeptide (ANSP) confirmed

that this band is the cacao trypsin inhibitor protein (accession X56509). In order to verify whether the endoproteinase was effectively composed of two subunits (29 and 13 kDa) (Voigt et al., J. Plant Physiol. 145 (1995), 299-307), several polypeptides smaller than 15.6 kDa were also sequenced. All the examined bands were found to be fragments of the 20.2 kDa cacao trypsin 5 inhibitor protein and none corresponded to a putative 13 kDa of TcAP2. Furthermore, the fact that the 30.5 kDa polypeptide contains both catalytic triads (D¹⁰⁸TG, D²⁹⁵SG) supports the idea that this polypeptide alone is proteolytically active. Therefore, TcAP2 is a novel monomeric aspartic endoproteinase.

10 **Example 7**

Characterisation of the native purified aspartic endoproteinase activity

Inhibitor Sensitivity: The inhibitor sensitivity of the native aspartic endoproteinase was determined in 300 µl reactions containing 200 mM sodium citrate, pH 3, 10 mg/ml bovine 15 hemoglobin, and 5 µl of size exclusion purified pool #57-64 (2.4 µg protein/µl). The inhibitors were added to give a final concentration of 2 µM pepstatin, 2 mM 1,10 phenanthroline, 100 µM dichloroisocoumarin (DCI), 10 µM E-64. The enzyme activity was determined as described in example 5. The fact that only pepstatin A inhibits completely the activity (Table 2) confirms that the protease activity purified is an aspartic endoproteinase.

20

Table 2 Inhibitor sensitivity of the purified aspartic endoproteinase activity. Two replicates were done for each test.

Inhibitor	mM	Remaining Activity %
-	-	100
Pepstatin A	0.002	0%
1,10 Phenanthroline	2.0	86%
E-64	0.01	88%
DCI	0.1	90%

25

Determination of the optimum pH: The activity test performed at different pH values indicated that the purified enzyme had an optimal activity at pH 3.0 (data not shown).

Example 8

Analysis of the products formed when a partially purified aspartic endoproteinase preparation is incubated in acid conditions

5

To examine the peptides produced by the native cacao seed aspartic endoproteinase, a Q Sepharose Fast Flow partially purified preparation of TcAP2 (197 µg protein, 1.35 units of activity/µl; specific activity 821 units/mg protein) was incubated in acid conditions. 120 µl of the partially purified enzyme were mixed with 30 µl 1 M sodium citrate pH 3. Samples of 4 µl and

10 70 µl were taken out just before incubation at 42°C (t=1 min) and after seven hours. The 4 µl samples were put in SDS gel loading buffer for SDS-PAGE analysis. The reaction in the 70 µl samples was stopped by adding SDS to 1% final concentration, the samples were freeze-dried, solubilized with 100 µl 6M urea, 20 mM sodium phosphate pH 7, loaded on a Superdex Peptide HR 10/30 column (Amersham Pharmacia Biotech) and eluted with 6M urea, 20 mM sodium phosphate pH 7 at ambient temperature.

The gel presented in Fig. 14 shows that after 7 hours, nearly all the proteins seen in the 1 min sample were substantially hydrolysed. Only two significant bands remain, one of which corresponds to a reduced amount of the 30.5 kDa cacao aspartic endoproteinase polypeptide

20 indicating an enhanced resistance of the aspartic endoproteinase towards autocatalytic degradation. When the products of the aspartic endoproteinase digestion were examined by high resolution size exclusion chromatography (Fig. 15), a significant proportion of small oligopeptides were detected, with a large percentage of the peptides having sizes ranging between 2 and 70 amino acids. This observation indicates that reacting the main cacao seed 25 aspartic endoproteinase (TcAP2) with proteins can generate a significant level of very small peptides, and thus that the action of this enzyme could generate a significant proportion of the cocoa flavor precursor peptides found in fermented cocoa beans.

30

Claims

1. A recombinant aspartic endoproteinase as identified by SEQ ID No. 1 or a variant thereof obtained by substituting, deleting or adding one or more amino acids with the proviso that the enzymatic activity of the aspartic endoproteinase is essentially retained.
2. A recombinant aspartic endoproteinase as identified by SEQ ID No. 2 or a variant thereof obtained by substituting, deleting or adding one or more amino acids with the proviso that the enzymatic activity of the aspartic endoproteinase is essentially retained.
3. A DNA sequence coding for an aspartic endoproteinase according to claim 1.
4. The DNA sequence according to claim 3, which is identified by SEQ ID No. 3 or a variant thereof obtained by replacing, deleting or adding one or more nucleotides such that the polypeptide coded thereby is still essentially active.
5. A DNA sequence coding for an aspartic endoproteinase according to claim 2.
6. The DNA sequence according to claim 5, which is identified by SEQ ID No. 4 or a variant thereof obtained by replacing, deleting or adding one or more nucleotides such that the polypeptide coded thereby is still essentially active.
7. A vector comprising a DNA sequence according to any of the claims 3 to 6.
8. A cell containing a recombinant DNA sequence according to any of the claims 3 to 7.
9. The cell according to claim 8, which is a prokaryotic cell, an eukaryotic cell or a plant cell, preferably a cacao cell.

10. A transgenic plant, containing a cell according to claim 9.
11. Use of a DNA sequence according to any of the claims 3 to 6 for the manufacture of a
5 cacao aspartic endoproteinase.
12. The use according to claim 11, wherein the aspartic endoproteinase is produced in a
suitable cell.
- 10 13. The use according to claim 12, wherein the cell is a prokaryotic or eukaryotic host cell
or a plant cell, preferably a cacao plant cell.
14. Use of an aspartic endoproteinase according to any of the claims 1 or 2 for the
preparation of cocoa flavour.
15
15. Use of an aspartic endoproteinase according to any of the claims 1 or 2 for
hydrolyzing proteins.
16. The use according to claim 15, wherein the proteins are derived from food material.
20
17. A process for producing cocoa flavour comprising subjecting a material suitable to
yield cocoa flavour precursors to an enzymatic degradation, involving the use of an
aspartic endoproteinase according to any of the claims 1 or 2.
- 25 18. A product containing cocoa flavour, obtainable according to the method of claim 17.
19. A process of hydrolyzing proteinaceous material in a plant comprising expressing an
aspartic endoproteinase according to any of claim 1 or 2 in plant cells, especially seed
cells, and then effecting hydrolysis of the cellular protein by treating such plant cells
30 with an acidic solution.

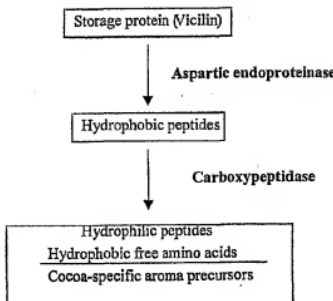


Fig. 1: Proteolytic formation of the cocoa-specific aroma

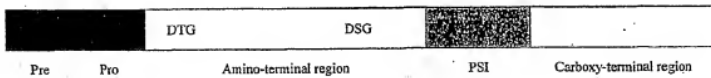


Fig. 2: Schematic representation of plant aspartic prepropeptides.

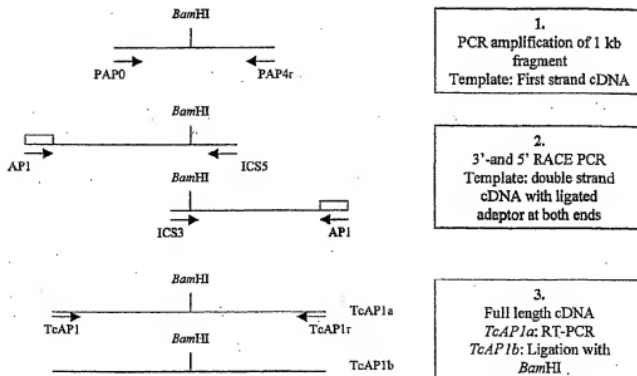


Fig. 3: Cloning strategy used for the isolation of cDNA encoding aspartic endoproteinase from *Theobroma cacao*.

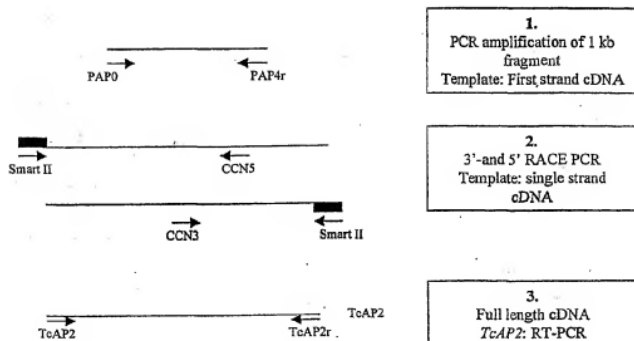


Fig. 4: Cloning strategy used for the isolation of cDNA encoding aspartic endoproteinase from *Theobroma cacao*,

	10	20	30	40	50
TcAP1a	SGRIVKTTT	TVLFLCLL	FFLPIVFSIS	NEKLRVRLK	KRKPDQNYRLA
TcAP1b	SGRIVKTTT	TVLFLCLL	FFLPIVFSIS	NEKLRVRLK	KRKPDQNYRLA
TcAP2	SGTTI	KKVVVLSL	LISSLLFPIVFS	SSVSDG	VLKIGLEEMELDQ
	60	70	80	90	100
TcAP1a	DSKEREAFR	ASLKKYRLQG	NLGESEDID	IVALKNRYD	DAQYFGEIGIG
TcAP1b	DSKEREAFR	ASLKKYRLQG	NLGESEDID	IVALKNRYD	DAQYFGEIGIG
TcAP2	DSHDG	EADRAFIKKYR	FRBMLG	DSEED	IVALKNRYD
	110	120	130	140	150
TcAP1a	QHFTV	I	DTGSSNLUV	SSKCYF	IACYLVHSRYKE
TcAP1b	QHFTV	I	DTGSSNLUV	SSKCYF	IACYLVHSRYKE
TcAP2	QHFTV	I	DTGSSNLUV	S	TCYFSSVACVF
	160	170	180	190	200
TcAP1a	QYGTGAI	SQFFF	SDRVHQ	QFEP	IEATR
TcAP1b	QYGTGAI	SQFFF	SDRVHQ	QFEP	IEATR
TcAP2	QYGTGAI	SQFFF	SDRVHQ	QFEP	IEATR
	210	220	230	240	250
TcAP1a	GFOEISVGN	AVPVVLYN	WVH	QG	LFV
TcAP1b	GFOEISVGN	AVPVVLYN	WVH	QG	LFV
TcAP2	GFOEISVGN	AVPVVLYN	WVH	QG	LFV
	260	270	280	290	300
TcAP1a	DKHFKG	QDHTTYV	P	TRK	G
TcAP1b	DKHFKG	QDHTTYV	P	TRK	G
TcAP2	DKHFKG	QDHTTYV	P	TRK	G
	310	320	330	340	350
TcAP1a	ITGPTTA	I	ACVHNAIGA	S	QV
TcAP1b	ITGPTTA	I	ACVHNAIGA	S	QV
TcAP2	LAGFS	TW	TM	I	ACVHNAIGA
	360	370	380	390	400
TcAP1a	SQIGL	C	DGTRGV	STG	IEB
TcAP1b	SQIGL	C	DGTRGV	STG	IEB
TcAP2	SQIGL	C	DGTRGV	STG	IEB
	410	420	430	440	450
TcAP1a	OLKQH	QSERL	IVINEL	CORL	PF
TcAP1b	OLKQH	QSERL	IVINEL	CORL	PF
TcAP2	OLKQH	QSERL	IVINEL	CORL	PF
	460	470	480	490	500
TcAP1a	FE	LSFEGV	V	LDV	PPR
TcAP1b	FE	LSFEGV	V	LDV	PPR
TcAP2	FDLT	P	LSFEGV	V	LDV
	510				
TcAP1a	PDYGH	I	QVGF	AA	AA
TcAP1b	PDYGH	I	QVGF	AA	AA
TcAP2	PDYGH	I	QVGF	AA	AA

FIG. 5

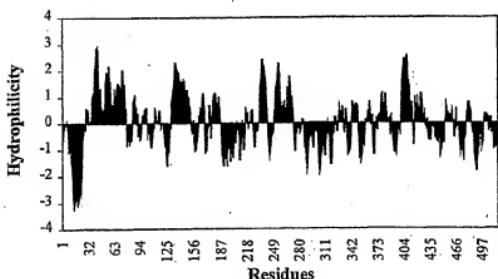
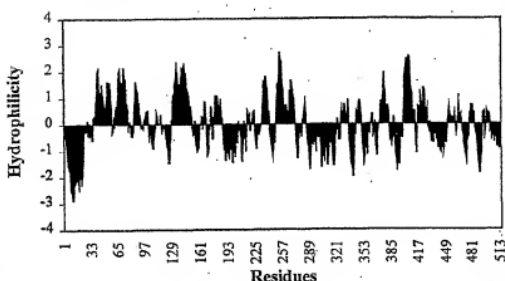
A**Hydrophilicity Plot-kyte-Doolittle for TcAP1a****B****Hydrophilicity Plot-kyte-Doolittle for TcAP2**

Fig. 6: Hydrophilicity Plot-Kyte-Doolittle for the TcAP1a (A) and TcAP2 (B) sequences.

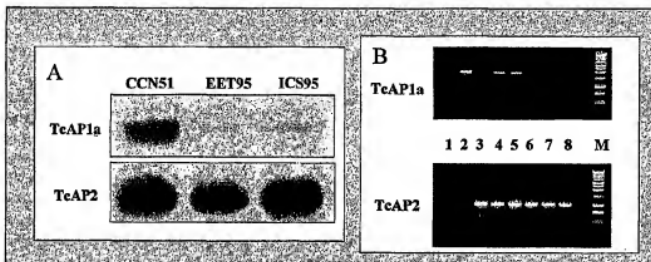


Fig. 7: *TcAP1a* and *TcAP2* expression in cacao bean produced by three different cacao clones

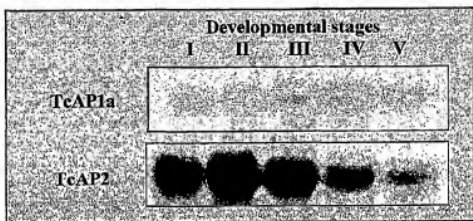


Fig. 8: Northern blot analysis of *TcAP1a* and *TcAP2*

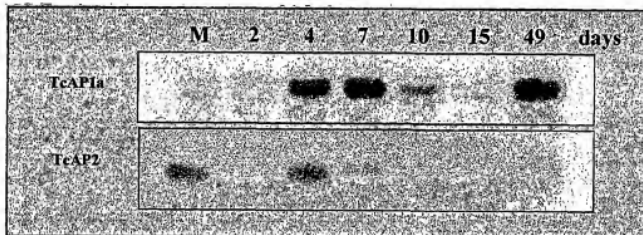


Fig. 9: Northern blot analysis of *TcAP1a* and *TcAP2* expression in cacao bean

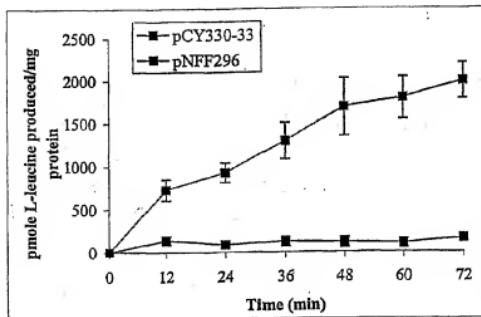


Figure 10: Hydrolysis of bovine haemoglobin by recombinant TcAP2 protein in yeast culture medium and comparison with control strain pNFF296.

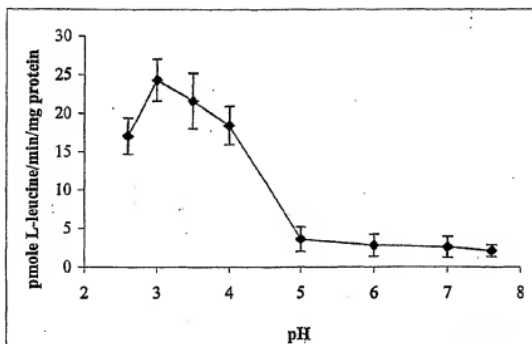


Figure 11: pH dependence of haemoglobin hydrolysis by recombinant TcAP2.

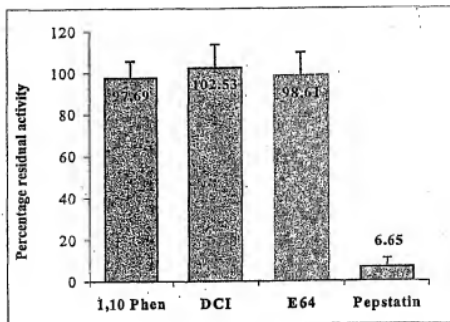


Figure 12: Effect of different inhibitors on the hydrolysis of bovine haemoglobin by recombinant TcAP2

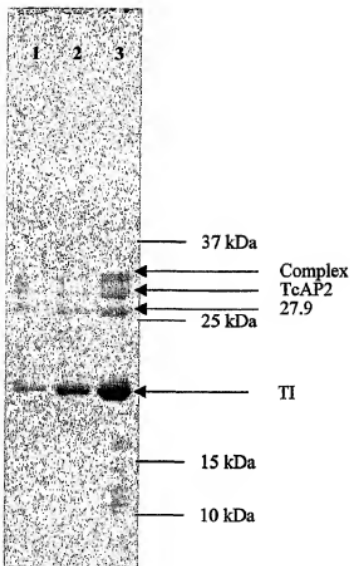


Fig. 13

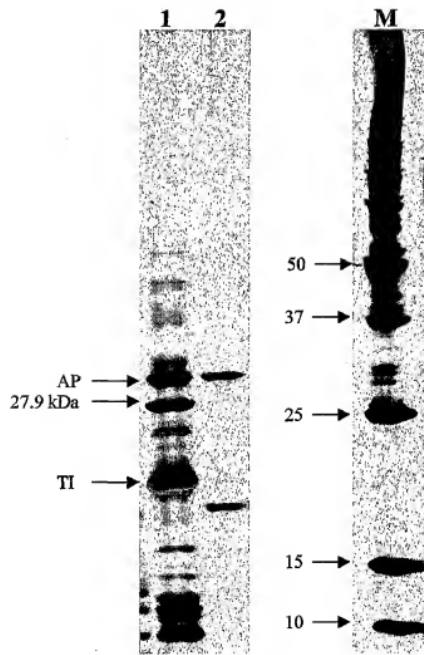


Fig. 14

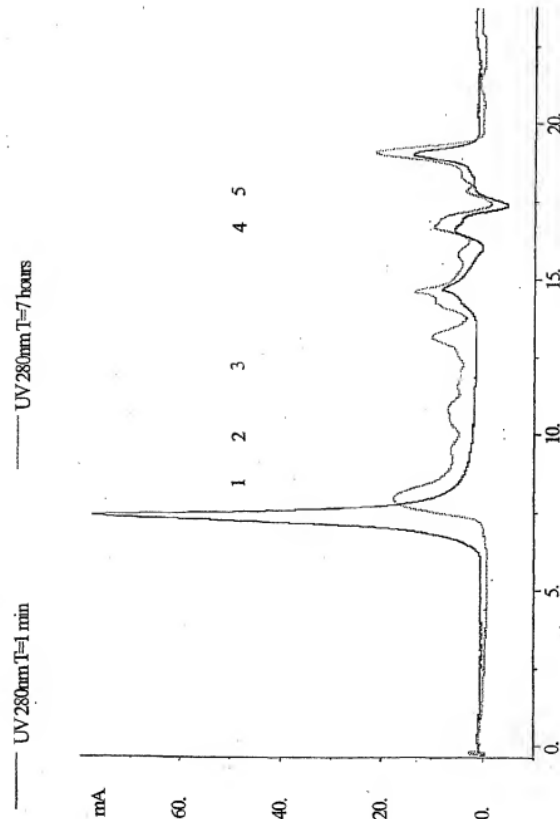


Fig. 15

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(74) Agent: STRAUS, Alexander; Becker, Kurig, Straus, Bavaristraße 7, 80336 München (DE).

(81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.

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WO 02/04617 A3

(54) Title: NOVEL CACAO ENDOPROTEINASES AND THEIR USE IN THE PRODUCTION OF COCOA FLAVOUR

(57) Abstract: The present invention pertains to novel aspartic endoproteinases from *Th. cacao* which are involved in the production of cocoa flavour and DNA sequences coding for them. In particular, the present invention relates to the use of said enzymes in the manufacture of cocoa flavour.

INTERNATIONAL SEARCH REPORT

International Application No
PCT/EP 01/07255A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N15/57 C12N9/50 A01H5/00 C12P21/06 A23G1/02

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 7 C12N A23G A01H C12P

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

BIOSIS, WPI Data, SEQUENCE SEARCH

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>VOIGT J ET AL: "Developmental stage-dependent variation of globular storage protein and aspartic endoprotease during ripening and germination of <i>Theobroma cacao</i> L. seeds." <i>JOURNAL OF PLANT PHYSIOLOGY</i>, vol. 145, no. 3, 1994, pages 299-307, XP000972681 ISSN: 0176-1617 page 300, right-hand column, paragraph 2</p> <p>---</p> <p>-/-</p>	1,2

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- *L* document which may throw doubt on priority, claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
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- *P* document published prior to the international filing date but later than the priority date claimed

T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

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INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/07255

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	<p>HIRAIWA NAGAKO ET AL: "An aspartic endopeptidase is involved in the breakdown of propeptides of storage proteins in protein-storage vacuoles of plants." EUROPEAN JOURNAL OF BIOCHEMISTRY, vol. 246, no. 1, 1997, pages 133-141, XP000971170 ISSN: 0014-2956</p> <p>-----</p>	
A	<p>EMBL DATABASE Accession no U61396, Sequence identity VUU61396 17 July 1996</p> <p>D'ARCY-LAMETA A ET AL: "Molecular cloning and nucleotide sequence of a cDNA encoding an aspartic proteinase from <i>Vigna unguiculata</i> leaves" XP002155324</p> <p>-----</p>	
A	<p>DATABASE BIOSIS 'Online!' BIOSCIENCES INFORMATION SERVICE, PHILADELPHIA, PA, US; 1995</p> <p>VOIGT J ET AL: "Precursors of the cocoa-specific aroma components are derived from the vicilin-class (7S) globulin of the cocoa seeds by proteolytic processing." Database accession no. PREV199698592431 XP002194847 abstract & BOTANICA ACTA, vol. 108, no. 4, 1995, pages 283-289, ISSN: 0932-8629</p> <p>-----</p>	14-19